SECTION 2

EPA TRACEABILITY PROTOCOL FOR ASSAY AND CERTIFICATION OF COMPRESSED GAS CALIBRATION STANDARDS

2.1 GENERAL INFORMATION

2.1.1 Purpose and Scope of the Protocol

This protocol describes two procedures for assaying the concentration of compressed gas calibration standards and for certifying that the assayed concentrations are traceable to a National Institute of Standards and Technology (NIST) Standard Reference Material (SRM). This protocol is mandatory for certifying the compressed gas calibration standards used for the pollutant monitoring that is required by the regulations of 40 CFR Parts 50, 58, 60 and 75 for the calibration and audit of ambient air quality analyzers and continuous emission monitors. This protocol may be used to assay and certify gas mixtures that have the same components as compressed gas SRMs. A multiple-component standard may be assayed and certified under this protocol if compressed gas SRMs that contain the individual components in the standard exist. This protocol may be used by specialty gas producers, standard users, or other analytical laboratories. The assay procedure may involve the direct comparison of the standards to reference standards without dilution (i.e., Procedure G1) or the indirect comparison of the standards to reference standards with dilution (i.e., Procedure G2). A candidate standard having a concentration that is lower or higher than that of the reference standard may be certified under this protocol if both concentrations (or diluted concentrations) fall within the well-characterized region of the pollution gas analyzer's calibration curve. This protocol places no restrictions on cylinder sizes and the same analytical procedures must be used in assays of all cylinder sizes.

2.1.2 Reference Standards

Parts 50, 58, 60, and 75 of the monitoring regulations require that gaseous pollutant concentration standards used for calibration and audit of ambient air quality analyzers and continuous emission monitors be traceable to either a NIST SRM or a NIST Traceable Reference Material (NTRM). In 1996, NIST and the Netherlands Measurement Institute (NMi) issued a joint declaration that specific NMi Primary Reference Materials (PRMs) can be considered as being equivalent to the corresponding NIST SRMs. The compressed gas SRMs that are available from NIST are listed in Table 2-1. The current SRM-equivalent compressed gas PRMs that are available from NMi are listed in Table 2-2. Other gas mixtures are under study by NIST and NMi and they may be added to the Declaration of Equivalence. PRMs produced by other national metrology organizations will be considered equivalent to NIST SRMs when a declaration of equivalence is issued jointly by NIST and the national metrology organization. The generic terms "Primary Reference Material" and "PRM" are used in this document to refer to any SRM-equivalent standard that has received such equivalency status.

The uncertainty of SRMs, NTRMs, and PRMs is expressed as a 95-percent confidence interval, which is the one-sigma uncertainty multiplied by a coverage factor almost always equal to

TABLE 2-1. SUMMARY OF COMPRESSED GAS SRMs
THAT ARE AVAILABLE FROM NIST^a

	Balance gas	Concentration ^b range
Certified component	Balarice yas	for SRMs
Ambient nonmethane organics (15 components)	Nitrogen	5 ppb
Ambient toxic organics (19 components)	Nitrogen	5 ppb
Aromatic organic gases ^c	Nitrogen	0.25 to 10 ppm
Carbon dioxide	Air	345 to 365 ppm
Carbon dioxide	Nitrogen	0.5 to 16 percent
Carbon monoxide	Air	10 to 45 ppm
Carbon monoxide	Nitrogen	10 ppm to 13 percent
Carbon monoxide, propane, and carbon dioxide	Nitrogen	1.6 to 8 percent CO 600 to 3,000 ppm C_3H_8 0 to 14 percent CO_2
Hydrogen sulfide	Nitrogen	5 to 20 ppm
Methane	Air	1 to 10 ppm
Methane	Nitrogen	50 to 100 ppm
Methane and propane	Air	4 ppm CH ₄ , 1 ppm C ₃ H ₈
Nitric oxide	Nitrogen	5 to 3,000 ppm
Oxides of nitrogen (i.e., nitrogen dioxide plus nitric acid)	Air	100 ppm
Oxygen	Nitrogen	2 to 21 percent
Propane	Air	0.25 to 500 ppm
Propane	Nitrogen	100 ppm to 2 percent
Sulfur dioxide	Nitrogen	50 to 3,500 ppm

^a All SRMs may not be available at all times. Other compressed gas SRMs may be developed in the future and could be used as reference standards. Contact NIST for information about SRM availability at (301) 975-6776 or http://gases.nist.gov.

b SRM concentrations are by mole.

^c Aromatic organic gases are benzene, bromobenzene, chlorobenzene, and toluene.

TABLE 2-2. SUMMARY OF COMPRESSED GAS PRMs
THAT ARE AVAILABLE FROM NMi^a

Certified component	Balance gas	Concentration range for PRMs ^b	
Carbon dioxide	Nitrogen	100 ppm to 15 percent	
Carbon monoxide	Nitrogen	100 ppm to 6 percent	
Ethanol	Nitrogen	100 to 259 ppm	
Nitric oxide	Nitrogen	10 to 4000 ppm	
Oxygen	Nitrogen	2 to 22 percent	
Propane	Nitrogen	500 to 3000 ppm	
Sulfur dioxide	Nitrogen	100 to 3500 ppm	

^a Information about PRMs can be obtained from:

The NMi office in the United States is:

Nederlands Meetinstituut B.V. Van Swinden Laboratorium Department of Chemistry P.O. Box 65\(P.O. Box 7758 2600 AR DELFT

The Netherlands
Telephone: 31 15 269 16 80
Telephone: 31 15 261 30 71

Telefax: 31 15 261 29 71 E-mail: SecChemie@NMi.nl NMi USA, Inc.

36 Gilbert Street South Tinton Falls, NJ 07701

Shrewsbury, NJ 07701 Telephone: (908) 842-8900 Telefax: (908) 842-0304 E-mail: NMiUSANJ@aol.com

SRM-equivalent PRMs from other national metrology organizations may be added in the future. Users of this protocol will be advised if such additions occur.

2.9 This estimate includes allowances for the uncertainties of known sources of systematic error as well as the random error of measurement. A value of one-half of the stated uncertainty of these reference standards should be used in calculating the total analytical uncertainty of standards that are certified under this protocol (see Appendix C).

The EPA regulations define a "traceable" standard as one that has been compared and certified, either directly or via not more than one intermediate standard, to a primary standard such as a NIST SRM or an NTRM. Comparison of a candidate standard directly to an SRM, an SRM-equivalent PRM, or an NTRM is preferred and recommended. However, the use of a Gas Manufacturer's Intermediate Standard (GMIS) (see Subsection 2.1.2.1) in the comparison is permitted. A GMIS is an intermediate reference standard that has been compared directly to an SRM, an SRM-equivalent PRM, or a NTRM according to Procedure G1. It is an acceptable reference standard for the assay of candidate standards. However, purchasers of standards that have been compared to a GMIS should be aware that, in conformity with the above definition, such a standard

Within the listed ranges, any concentration is available. PRMs are prepared individually in 5-L cylinders according to ISO Standard 6142 (Gas Analysis–Preparation of calibration gas mixtures-weighing methods). After preparation, the composition is verified against Dutch Primary Standard Gas Mixtures. The stability is normally guaranteed for a period of 2 years. Uncertainties depend on the certified concentration and vary from 0.1 percent (relative) for binary mixtures to 1.0 percent (relative) maximum for certain constituents in multicomponent mixtures.

could only be used directly for calibration or audit. Such a standard could not be used as a second-generation intermediate reference standard to certify other compressed gas calibration standards.

Accordingly, the reference standard used for assaying and certifying a compressed gas calibration standard under this protocol must be an SRM, a NTRM, an SRM-equivalent PRM, or a GMIS. The reference standard must be within its certification period.

Volume reference standards must be traceable to NIST primary standards by calibration at a NIST-accredited state weights and measures laboratory or at a calibration laboratory that is accredited by the National Voluntary Laboratory Accreditation Program (NVLAP) or by the International Laboratory Accreditation Conference (ILAC). These volume reference standards are required for assays using procedure G2 (see Subsection 2.3.7).

2.1.2.1 Gas Manufacturer's Intermediate Standard—

A GMIS is a compressed gas calibration standard that has been assayed by direct comparison to an SRM, an SRM-equivalent PRM, or a NTRM, that has been assayed and certified according to Procedure G1, and that also meets the following requirements:

- A candidate GMIS must be assayed on at least three separate dates that are uniformly spaced over at least a 3-month period. During each of these assays, the candidate GMIS must be measured at least three times. All these assays must use the same SRM, SRM-equivalent PRM, or NTRM as the reference standard to avoid errors associated with the use of different reference standards for different assays.
- 2. For each assay, the analyst must calculate the mean and 95-percent uncertainty for the three or more measured concentrations of the candidate GMIS according to the statistical procedures described in Appendix A or equivalent statistical techniques. The 95-percent uncertainty must be less than or equal to 1.0 percent of the mean concentration.
- 3. After the three or more assays have been completed, the analyst must calculate the overall mean estimated concentration and the 95-percent uncertainty for the candidate GMIS using the spreadsheet described in Appendix C or equivalent statistical techniques.
- 4. If the 95-percent confidence limits (i.e., estimated concentration plus or minus uncertainty) for the assays overlap, the candidate GMIS can be considered to be stable and can be used as a reference standard for assays of candidate standards. In the Appendix C spreadsheet, all cells in the comparisons table will be "true." If the confidence limits do not overlap, the candidate GMIS may be unstable or there may be analytical problems associated with the assays or the reference standards. One or more cells in the comparisons table will be "false." The analyst must either disqualify the candidate GMIS or investigate why the confidence limits do not overlap. The analyst may discard the data from a questionable assay and then conduct another assay. The candidate GMIS can be used as a reference standard if the confidence limits for the remaining assays and the new assay overlap. The candidate GMIS cannot be used if it appears to be unstable.
- 5. A GMIS must be recertified every 2 years. Use the spreadsheet described in Appendix C or equivalent statistical techniques to compare the confidence limits from a single recertification assay with the confidence limits from the previous assays. If the confidence intervals overlap, the GMIS can be recertified. If the reassayed GMIS fails to meet this requirement, it must undergo a full certification as described in Step 1 above before it can be used again. There is no requirement that the same reference standard must be used in the original assays and the recertification assay, but this practice is desirable if possible.

2.1.2.2 Recertification of Reference Standards—

Recertification requirements for SRMs and NTRMs are specified by NIST. Recertification requirements for PRMs are specified by NMi. See Subsection 2.1.2.1 for GMIS recertification requirements.

2.1.3 Using the Protocol

The assay/certification protocol described here is designed to minimize both systematic and random errors in the assay process. Therefore, the protocol should be carried out exactly as it is described. The assay procedures in this protocol include one or more possible designs for the assay apparatus. The analyst is not required to use these designs and may use alternative components and configurations that produce equivalent-quality measurements. Inert materials (e.g., Teflon[®], stainless steel, or glass) and clean, noncontaminating components should be used in those portions of the apparatus that are in contact with the gas mixtures being assayed.

2.1.4 Certification Documentation

Each certified compressed gas calibration standard must be documented in a written certification report and this report must contain at least the following information:

- 1. Cylinder identification number (e.g., stamped cylinder number).
- Certified concentration for the compressed gas calibration standard, in parts per million by mole or mole percent. This value should be reported to 3 significant digits. The certified concentration is the mean of all assayed concentrations for which the candidate standard is considered to be stable.
- 3. Balance gas in the gas mixture.
- 4. Cylinder pressure at certification and the statement that the standard should not be used when its gas pressure is below 1.0 megapascals (i.e., 150 psig).
- 5. Date of the assay/certification.
- 6. Certification expiration date (i.e., the certification date plus the certification period) (see Subsection 2.1.6.3).
- 7. Identification of the reference standard used in the assay: NIST SRM number, NIST sample number, cylinder identification number and certified concentration for an SRM; cylinder identification number and certified concentration for an SRM-equivalent PRM, a NTRM, or a GMIS. The certification documentation must identify the reference standard as being an SRM, an SRM-equivalent PRM, a NTRM, or a GMIS.
- 8. Statement that the assay/certification was performed according to this protocol and that lists the assay procedure (e.g., Procedure G1) used.
- 9. The analytical method that was used in the assay.
- 10. Identification of the specialty gas producer or other laboratory (i.e., name and location) where the standard was assayed and certified. This identification must be given in the same or larger font as the other required information in the report.
- 11. Chronological record of all certifications for the standard.

- 12. If applicable, statement that the certified concentration of specified component has been corrected for analytical interferences from other specified components.
- 13. An estimate of the total uncertainty associated with the assay of the candidate standard. This estimate must include the uncertainties of the reference standards, the analyzer multipoint calibration, and any interference correction. Use the spreadsheet described in Appendix C or equivalent statistical techniques to calculate the total uncertainty.

This certification documentation must be given to the purchaser of the standard. The specialty gas producer must maintain laboratory records and certification documentation for 3 years after the standard's certification date. A specialty gas producer or other vendor may redocument an assayed and certified standard that it has purchased from another specialty gas producer and that it wishes to sell to a third party. However, the new certification documentation must clearly identify the specialty gas producer or other laboratory (i.e., name and location) where the standard was assayed.

2.1.5 Certification Label

A label or tag bearing the information described in items 1-6, 8, and 10 of Subsection 2.1.4 must be attached to the standard.

2.1.6 Assay/Certification of Compressed Gas Calibration Standards

2.1.6.1 Incubation of Newly Prepared Compressed Gas Calibration Standards—

Newly prepared compressed gas calibration standards must be incubated at least 4 days before being assayed and certified.

2.1.6.2 Stability Test for Reactive Gas Mixtures—

Compressed gas calibration standards that contain reactive gas mixtures, including hydrogen sulfide (H₂S), nitric oxide (NO), oxides of nitrogen (NO), sulfur dioxide (SO), and carbon monoxide (CO), and that have not been previously certified, must be tested for stability as discussed herein. Conduct an initial assay of the candidate standard and determine a concentration for the standard. The candidate standard must be measured at least three times during the assay. Reassay the standard at least 7 days after the first assay. Use the spreadsheet described in Appendix C or equivalent statistical techniques to compare the 95-percent confidence limits for the two assays. If the confidence limits overlap, the candidate standard can be considered to be stable and can be certified. In the spreadsheet, all cells in the comparisons table will be "true." If the confidence intervals do not overlap, the candidate standard may be unstable or there may be analytical problems associated with the assays or the reference standards. The analyst must wait an additional 7 days or more and conduct a third assay. If the confidence interval for the third assay overlaps either of the two previous assays, the candidate standard can be certified using the data from the two overlapping assays to determine the certified concentration and the total uncertainty. The analyst must disqualify the candidate standard if none of the three confidence intervals overlap. The analyst should investigate the cause of the lack of agreement among the three assays and should correct any problems that are discovered.

2.1.6.3 Certification Periods for Compressed Gas Calibration Standards—

The certification of a compressed gas calibration standard is valid for only a specified period following its certification date, which is the date of its last assay. In general, the certification period should be no longer than the period for which similar standards have been shown to be stable. The certification periods for various standards are specified in Table 2-3. These certification periods are for standards that are contained in aluminum cylinders. In general, the certification period for standards that are contained in nonaluminum cylinders is 6 months. However, an exception is made for the following three gas mixtures: carbon dioxide with a concentration >0.5 percent; oxygen with a

concentration >0.5 percent; oxygen with a concentration >0.8 percent; and propane with a concentration >0.1 percent. The certification period for standards containing these three gas mixtures in nonaluminum cylinders is given in Table 2-3.

If a standard is to be used after its certification period has ended, it must be recertified in accordance with this protocol. The recertification assay must be performed using the same analytical procedure (e.g., Procedure G1) as was used for the original assay of the standard. The purpose of this assay is to determine whether the standard has remained stable since its original certification. The standard must be measured at least three times during the recertification assay. A second assay is not needed for recertification of the standard. There is no requirement that the same reference standard must be used in the original and recertification assays, although this practice is desirable if possible. Record the results of the recertification assay in the laboratory's records. Use the spreadsheet described in Appendix C or equivalent statistical techniques to compare the confidence limits for the recertification assay with those for the previous assays. If the confidence limits overlap, the standard can be recertified. The second certification period for the standard is the same as that given in Table 2-3.

A standard that was certified under this protocol may be recertified by a laboratory other than the one that performed the original certification. In such a case, the 95-percent confidence limits for the recertification assay must overlap the certified concentration plus or minus the total uncertainty that was given in the original certification documentation. If the confidence limits do not overlap, a second recertification assay must be conducted and the confidence limits for the two recertification assays must overlap before the standard can be recertified. The recertification documentation must list the information from the original certification documentation plus the corresponding information from the recertification assays. Both the original and the recertification laboratories must be identified in the recertification documentation.

TABLE 2-3. CERTIFICATION PERIODS FOR COMPRESSED GAS CALIBRATION STANDARDS IN ALUMINUM CYLINDERS THAT ARE CERTIFIED UNDER THIS PROTOCOL

Certified components	Balance gas	Applicable concentrationrange	Certification period (months)
Ambient nonmethane organics (15 components)	Nitrogen	5 ppb	24
Ambient toxic organics (19 components)	Nitrogen	5 ppb	24
Aromatic organic gases	Nitrogen	≥0.25 ppm	36
Carbon dioxide	Nitrogen or air ^a	≥300 ppm	36
Carbon monoxide	Nitrogen or air	≥8 ppm	36
Hydrogen sulfide	Nitrogen	≥4 ppm	12
Methane	Nitrogen or air	≥1 ppm	36
Nitric oxide	Oxygen-free nitrogen ^b	≥4 ppm	24
Nitrous oxide	Air	≥300 ppb	36
Oxides of nitrogen (i.e., sum of nitrogen dioxide and nitric acid)	Air	≥80 ppm	24
Oxygen	Nitrogen	≥0.8%	36
Propane	Nitrogen or air	≥1 ppm	36
Sulfur dioxide	Nitrogen or air	40 to 499 ppm	24
Sulfur dioxide	Nitrogen or air	≥500 ppm	36
Multicomponent mixtures	-	_	See text
Mixtures with lower concentrations	_	_	See text

^a When used as a balance gas, "air" is defined as a mixture of oxygen and nitrogen where the minimum concentration of oxygen is 10 percent and the concentration of nitrogen is greater than 60 percent.

The spreadsheet described in Appendix C to calculate the total analytical uncertainty of a candidate standard has provision for data from only three assays. If more than three assays are conducted, only the data from the three most recent assays should be used in the spreadsheet.

The certification periods given in Table 2-3 apply to specific concentration ranges over which the gas mixtures have been found to be stable. These concentration ranges match the concentration ranges for NIST SRMs. The protocol described here allows the certification of standards with concentrations that may be lower than those of the corresponding SRMs. If the concentration of the standard is less than the applicable concentration range given in Table 2-3, the initial certification period for this standard is 6 months. After this period, the standard must be recertified before further use. The standard must be measured at least three times during the recertification assay. If the confidence limits for the recertification assay overlap those for the previous assays, the standard can be recertified for the period shown in Table 2-3. For example, a 35-ppm sulfur dioxide in nitrogen standard will have an initial

^b Oxygen-free nitrogen contains <0.5 ppm of oxygen.

certification period of 6 months. After a successful recertification, this standard will have a recertification period of 24 months.

If the confidence limits from the recertification assay do not overlap those from the original assays, the analyst must either disqualify the standard for further use under this protocol or investigate why there is an apparent difference between the original assays and the recertification assay. This difference may be due to an actual instability of the gas mixture, to a reference standard problem, to an analytical instrumentation problem, or to some other problem. If the analyst can find a reasonable explanation for the difference and if this cause is not instability, then the standard can be recertified. The analyst must append a brief report on the investigation to the recertification documentation and to the laboratory's records.

A multiple-component standard can be certified for a period equal to that of its most briefly certifiable component. For example, a standard containing sulfur dioxide, carbon monoxide, and propane in nitrogen can be certified for 24 months because the shortest certification period is 24 months.

A standard may be recertified if the gas pressure remaining in the cylinder is greater than 3.4 megapascals (i.e., 500 psig).

2.1.6.4 Minimum Cylinder Pressure—

In general, a compressed gas calibration standard should not be used when its gas pressure is below 1.0 megapascals (i.e., 150 psig). NIST has found that some gas mixtures (e.g., nitric oxide in nitrogen) have exhibited a concentration change when the cylinder pressure fell below this value.

2.1.6.5 Assay/Certification of Multicomponent Compressed Gas Calibration Standards—

This protocol may be used to assay and certify a multiple-component standard if compressed gas SRMs, SRM-equivalent PRMs, or NTRMs exist that contain the individual components of the multiple-component standard. If any component in the multiple-component standard interferes with the assay of any other component, the analyst must conduct an interference study to determine an interference correction equation. This study must be conducted using the same analyzer or analyzers as will be used to assay the standard. The study must use single-component and multiple-component reference standards that have been assayed using interference-free analyzers. The study must cover the same range of concentrations for all components as will exist for the standards being assayed and certified according to this protocol.

Data from the interference study must be evaluated using multiple-variable least-squares regression analysis. The analyst should consult with a statistician before beginning the study or evaluating its data. The regression analysis must produce an interference correction equation and an estimate of the 95-percent uncertainty associated with the corrected concentrations for the assayed components. The interference correction equation will be valid for the range of concentrations covered in the study for which the uncertainty of the corrected concentration is ≤ 1 percent of the corrected concentration. The analyst must add the interference correction uncertainty to the total uncertainty of the standard. The certification documentation must include a statement that the certified concentration of a specified component has been corrected for interferences from other specified components. An interference study is not needed if the assay analyzer is interference free.

2.1.7 Analyzer Calibration

2.1.7.1 Basic Analyzer Calibration Requirements—

The assay procedures described in this protocol employ a data reduction technique to calculate the concentration of a candidate compressed gas standard that corrects for minor analyzer calibration variations (i.e., drift). This technique does not require the absolute accuracy of the analyzer's calibration curve at the time of the assay. Requirements for the analyzer follow: (1) it must have a well-

characterized calibration curve for the pollutant of interest (see Subsection 2.1.7.2); (2) it must have good resolution and low noise; (3) its calibration must be known and must be reasonably stable or recoverable during the assay/certification process; and (4) all measurements of candidate standards must fall within a well-characterized region of its calibration curve.

2.1.7.2 Analyzer Multipoint Calibration—

The gas analyzer used for the assay must have had a multipoint calibration within 1 month prior to the assay date. This calibration is not used directly to interpret analyzer response during the assay of the candidate compressed gas calibration standard. The data reduction technique corrects the analyzer response on the assay date for any minor calibration drift during the period between the multipoint calibration and the assay date. The corrected analyzer response is then used with the multipoint calibration data to calculate a concentration value for the candidate standard.

The analyzer's zero and span controls may be adjusted before the start of the multipoint calibration. If a zero or span adjustment is made, allow the analyzer to stabilize for at least one hour before beginning the multipoint calibration. The waiting period is necessary because some analyzers' calibrations drift for a period of time following a zero or span control adjustment.

The multipoint calibration must consist of one or more measurements of the analyzer responses to at least five different concentrations. The use of a zero gas in the calibration is recommended, but is not required. Record these measurements and the analyzer's zero and span control settings in the laboratory's records. These calibration concentrations should be approximately evenly spaced over the concentration range. The multipoint calibration is valid only for the concentration range lying between the largest and smallest measured concentrations. The concentrations may be produced by undiluted reference standards or by dilution of reference standards using a gas dilution system. See Subsection 2.1.7.4 for reference standard requirements. If a gas dilution system is used, it must have a specified accuracy of no worse than 1.0 percent of the undiluted reference standard concentration. The accuracy of the gas dilution system must be checked by the analyst at monthly intervals by comparing diluted reference standards to undiluted reference standards having approximately the same concentration.

If the analyzer has multiple concentration ranges, a multipoint calibration should be done for all ranges that will be used later for the assay of candidate standards. A multipoint calibration that is conducted on one range is not valid for an assay that is conducted on another range.

Data from the multipoint calibration must be evaluated using least-squares regression analysis. ¹⁵ This statistical analysis technique will be used to determine the analyzer's calibration curve and to characterize the uncertainty associated with the calibration. The concentration values are the independent (i.e., X) values in the analysis and their units may be parts per million, mole percent, or any other appropriate units. The analyzer response values are the dependent (i.e., Y) values in the analysis and their units may be volts, millivolts, percent of scale or any other measurable analyzer response units. The analyzer response values should have a resolution of less than or equal to 1 percent of the maximum measured analyzer response.

Because an analyzer's response has a random error component, repeated measurements of the same reference standard will not produce identical analyzer responses. The analyst may investigate the analyzer's precision by making replicate measurements of multiple reference standards. Least-squares regression analysis is normally conducted under the assumption that the precision is the same at all concentrations. However, this assumption may not be true for real-world analyzers and the analyst may need to use alternate statistical procedures to analyze the multipoint calibration data.

Calculate the least-squares regression coefficients of the calibration equation using the spreadsheets described in Appendix A or using equivalent statistical techniques (e.g., the worksheet for

linear relationships given in Chapter 5 of Reference 15). The spreadsheets allow the multipoint calibration data to be fitted to straight-line, quadratic, cubic, or quartic linear regression models. EPA discourages the use of the cubic and quartic models and believes that better fits of the data can be obtained by performing multipoint calibrations over more limited concentration ranges and by using straight-line or quadratic models. Inclusion of cubic and quartic models in the spreadsheets is for experimental use or for situations in which there is a theoretical basis for the use of such higher-order models. Analysts should be aware that apparent higher-order calibration curves may be caused by artifacts such as inaccurate reference standards or leaks in a gas dilution system. They should not use higher-order regression models to fit multipoint calibration data that have inadequate precision and that should be fitted to lower-order regression models. If analysts suspect that the precision is inadequate, they should make replicate measurements at each different concentration. Additionally, a multipoint calibration should not change regression model orders from one month to the next.

The spreadsheet described in Appendix A will suggest the best regression model for the multipoint calibration data, but the analyst should choose the model that best fits the measurement process on theoretical grounds.

Plot the values from the multipoint calibration and the regression curve with confidence bands as shown in Figure 2-1. These plots will provide a graphical representation of the calibration and will permit a qualitative assessment of the uncertainty associated with the calibration. Record the regression coefficients and other statistical results in the laboratory's records.

However, a quantitative assessment of the calibration's uncertainty is needed to allow the analyst to determine whether the multipoint calibration data adequately characterizes the "true" calibration curve for the analyzer. The criterion to be used to evaluate the uncertainty of the multipoint calibration is the 95-percent uncertainty (i.e., $\alpha = 0.05$) for a concentration predicted from the regression line using measured values of the analyzer response. This 95-percent uncertainty for the predicted concentration can be calculated using the spreadsheets described in Appendix A or using equivalent statistical techniques. Record the uncertainty calculations in the laboratory's records. A multipoint calibration will be considered to be well-characterized for all concentrations that are within the range of the multipoint calibration measurements and for which the magnitude of the 95-percent confidence limits for the regression-predicted analyzer response are $\leq \pm 1$ percent of the measured response for the largest concentration in the multipoint calibration. For example, assume that a multipoint calibration was conducted between 0 and 100 ppm and that the measured responses ranged between 0 and 10 volts. The calibration is well-characterized for all concentrations for which the 95-percent confidence limits are $\leq \pm 0.1$ volt, which is equal to ± 1 percent of 10 volts. Step 4 of the spreadsheet described in Appendix A allows the analyst to enter various concentrations and obtain the corresponding regression-

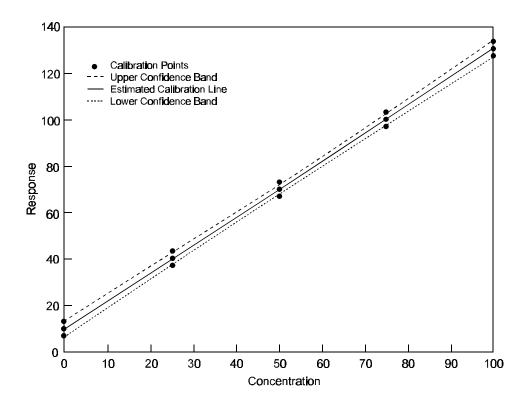


Figure 2-1. Example regression curve and confidence bands from multipoint calibration.

predicted analyzer response and confidence limits.

In effect, the 95-percent uncertainty value is a measure of how well the multipoint calibration data fit an equation which the analyst assumes is the "true" calibration equation for the analyzer. Comparison of uncertainty values from straight-line and quadratic equations permits the analyst to select the equation that best represents the calibration data.

A multipoint calibration may fail to meet this uncertainty criterion for several possible reasons:

- inadequate analytical precision;
- inaccuracy of the reference standards or the gas dilution system; or
- excessive uncertainty in the analyzer's calibration equation due to incorrect assumptions about the form of the equation.

The effect of inadequate analytical precision can be reduced by increasing the number of replicate measurements at each calibration concentration or by increasing the number of different concentrations used in the multipoint calibration. Additionally, precision can be improved by using an averaged analyzer response, rather than an instantaneous analyzer response, for each measurement. Reference standard inaccuracy is reduced by using the most accurate reference standards that are available. An inaccurate gas dilution system can be detected by comparing measurements of the concentration of a diluted reference

standard to the theoretically equal concentration of another, undiluted reference standard. It can also be detected by comparing measurements of two theoretically equal concentrations obtained by dilution of two reference standards having significantly different concentrations. An inaccurate gas dilution system must not be used for the multipoint calibration. The effect of excessive uncertainty in a straight-line calibration equation can be eliminated by using a quadratic calibration equation or by transforming the calibration data mathematically so that they may be fitted to a straight line regression equation. See Subsection 2.1.7.5 for a discussion of such mathematical transformations.

Note that possibly a more restrictive uncertainty criterion applies for the assay of the candidate standard. The magnitude of the 95-percent confidence limits for the estimated concentration of the candidate standard must be $\le \pm 1$ percent of the concentration of the reference standard (see Subsections 2.2.2 and 2.32). For example, assume that a 70-ppm candidate standard is being assayed using a 50-ppm reference standard. The 95-percent confidence limits for the candidate standard's estimated concentration must be $\le \pm 0.5$ ppm.

2.1.7.3 Zero and Span Gas Checks—

On any day after the multipoint calibration that the analyzer will be used for the assay of a candidate standard, its calibration drift must be measured. This drift is calculated relative to the analyzer response during the multipoint calibration. The purpose of the zero and span gas checks is to verify that the calibration drift has remained within acceptable limits since the multipoint calibration. The criterion that is used to assess the drift is the relative difference between the analyzer's current response and the corresponding value from the multipoint calibration. The following equation is used for this calculation:

Relative Difference =
$$100 \left[\frac{\text{Current Response - Calibration Response}}{\text{Calibration Response for Reference Standard}} \right]$$
.

This calculation is performed in Step 6 of the spreadsheet described in Appendix A.

Note that the relative difference is always calculated relative to the calibration response for the reference standard, even when the zero gas is being measured. This calculation is performed for the zero gas measurements and for the reference standard measurements. If the reference standard was not measured during the multipoint calibration, use the regression-predicted response for a concentration equal to that of the reference standard.

If the relative differences for the zero and span gas checks are each less than or equal to 5.0 percent, the analyzer's current calibration is considered to be approximately the same as during the multipoint calibration and the assay may be conducted. The zero and span controls do not have to be adjusted following the zero and span checks because the data reduction technique used in this protocol does not depend on the absolute accuracy of the analyzer calibration equation at the time of the assay.

If the relative differences for the zero or span gas checks are greater than 5.0 percent, the analyzer is considered to be out of calibration. A new multipoint calibration may be conducted before the candidate standard is assayed or the analyzer's zero and span controls may be adjusted to return the analyzer's response to the original calibration levels. For some analyzers such as nondispersive infrared instruments, daily changes in environmental variables such as barometric pressure may shift the calibration. After any adjustment of controls, the analyst should repeat the zero and span gas checks and recalculate the relative differences to verify that the analyzer is in calibration.

The zero gas and reference standard measurements that are performed for the assay of the candidate standard may also be used for the zero and span gas checks.

Between the time of the multipoint calibration and the time of the zero and span gas checks, the analyst may adjust the analyzer's zero and span controls for assays that will not be certified according to this protocol. However, these controls must be returned to their settings at the multipoint calibration before the zero and span gas checks or assays under this protocol.

2.1.7.4 Reference Standards for Multipoint Calibrations and Zero and Span Gas Checks—

The reference standards for the multipoint calibration and for the span gas checks must be diluted or undiluted SRMs, SRM-equivalent PRMs, NTRMs, or GMISs as specified in Subsection 2.1.2. The reference standard for the span gas check need not be the same as one of those used for the multipoint calibration or for the assay of the candidate standard.

Pure gases may be diluted to prepare gas mixtures for use in multipoint calibrations, but such mixtures may not be used as the reference standards for the span gas check or for the assay of the candidate standard. Pure gases may not be diluted by more than a factor of 100.

The zero gas must meet the requirements in Subsection 2.1.9. For some analyzers such as gas chromatographs, the analyst may have reason to believe that the zero gas reading may not accurately represent the zero-intercept of the calibration equation. The analyst may substitute a low-concentration, NIST-traceable reference standard for the zero gas, providing that the concentration of this standard is less than the concentration of the candidate standard.

2.1.7.5 Uncertainty of the Calibration Curve—

The data reduction technique used in this protocol is based on the assumption that the analyzer has a well-characterized calibration curve. The accuracy of the certified concentration of a candidate compressed gas calibration standard is dependent upon this assumption. The analyst cannot assume that the analyzer's calibration curve is a straight line between the measured values for the zero gas and the reference standard. The analyst must calculate the calibration equation and the uncertainty for its predicted concentrations by statistical analysis of the measurements obtained during the multipoint calibration.

The total uncertainty of the certified concentration for a candidate standard is composed of several components. The first component is the accuracy associated with the certified concentration of the reference standard. This uncertainty is minimized by using an SRM, an SRM-equivalent PRM, a NTRM, or a GMIS as the reference standard. The second component is the precision of the measurements of the reference and candidate standards. This uncertainty is minimized by making replicate measurements of these standards. The third component is the uncertainty associated with the concentrations that are predicted from the analyzer's calibration curve. This uncertainty concerns whether an assumed calibration equation accurately represents the "true" calibration curve.

This third component of uncertainty does not exist if the concentrations of the reference and candidate standards are equal. The assumed calibration equation and the true calibration curve will pass through the data for the reference standard regardless of whether they diverge elsewhere and the equation will be accurate for that single concentration. However, the uncertainty does exist if the concentrations of the reference and candidate standards differ. The assumed and true calibration curves may pass through different points for concentrations not equal to that of the reference standard. Analytical errors will develop because of this difference.

The measure of this uncertainty that is most directly useful to the analyst is the 95-percent uncertainty for a regression-predicted concentration given one or more measurements of the candidate

standard. The uncertainty may be calculated using the spreadsheet described in Appendix A or using equivalent statistical techniques. Several points should be noted about this uncertainty value. First, its magnitude decreases as n increases where n is the number of measurements in the multipoint calibration. Second, its magnitude decreases as n' increases, where n' is the number of measurements of the candidate standard. Third, its magnitude increases as the mean measured analyzer response ($\frac{1}{3}$) for the candidate standard diverges from the overall mean measured analyzer response ($\frac{1}{3}$) for the multipoint calibration. These points mean that it becomes easier to satisfy the uncertainty criterion as one increases the number of measurements in the multipoint calibration and in the assay of the candidate standard. Additionally, the absolute uncertainty of the regression predicted concentration is larger at the extremes of the calibrated concentration range than at the middle of the range.

For analyzers having an inherently nonlinear, but precise response, the calibration equation can be calculated using quadratic or higher-order polynomial regression analysis. Alternatively, a nonlinear equation may be linearized with a simple mathematical transformation of the multipoint calibration data. Examples of some linearizing transformations are given in Table 2-4, which is reproduced from Reference 15. The multipoint calibration data may need to undergo several different transformations before the optimum transformation is determined. Using appropriately transformed calibration data, a calibration equation can be calculated with an acceptable 95-percent uncertainty for the regression-predicted concentration. Subsequently, data obtained from the assay of the candidate standard must be similarly transformed to calculate a concentration for the candidate standard.

2.1.8 Uncertainty of the Estimated Concentration of the Candidate Standard

Uncertainty in the concentration estimated for a candidate standard is due to many different sources, including uncertainty in the reference standards, uncertainty in the analyzer multipoint calibration, uncertainty in the zero/span correction factors, and measurement imprecision. Some of these sources can be assessed using standard statistical techniques, but others cannot be assessed with the limited data that are produced when implementing this protocol.

For those cases where the candidate standard is assayed at the same time as the multipoint calibration, the candidate standard's concentration is determined directly from the calibration curve. The total uncertainty of the concentration is calculated by using the spreadsheets described in

TABLE 2-4. SOME LINEARIZING TRANSFORMATIONS FOR MULTIPOINT CALIBRATION DATA

If the relationship is of the form:	STEP ONE: Plot the transformed calibration data		STEP TWO: Fit the straight line	STEP THREE: Convert straight line constants (b ₀ and b ₁) to original constants	
	$Y_T =$	$X_T =$	$Y_T = b_0 + b_1 X_T$	b ₀ =	b ₁ =
$Y = a + \frac{b}{X}$	Υ	$\frac{1}{X}$	Use the normal procedures for calculating the regression line using	а	b
$Y = \frac{1}{a + bX'},$ or $\frac{1}{X} = a + bX$	1 Y	X	the transformed calibration data. Calculate the 95-percent uncertainties for the predicted transformed concentrations and	а	b
$Y = \frac{X}{a + bX}$	$\frac{X}{Y}$	X	compare them to the uncertainty criterion.	a	b
$Y = ab^x$	log Y	X		log a	log b
$Y = ae^{bx}$	log Y	Х		log a	b log e
aX ^b	log Y	log X		log a	b
Y = a + bX ⁿ , where n is known	Υ	X ⁿ		а	b

Source: Reference 15.

Appendices A and C or equivalent statistical techniques. It combines the uncertainty of the assay with the uncertainty of the reference standard using the following equation:

Uncertainty_{TOTAL} =
$$\sqrt{\text{(Uncertainty}_{ASSAY})^2 + \text{(Uncertainty}_{STANDARD})^2}$$
.

For those cases where the candidate standard is assayed on a date following the multipoint calibration, the spreadsheet includes the uncertainty associated with the zero gas and reference standard measurements in the calculation of total uncertainty.

If an interference-correction equation has been used to obtain a corrected concentration for the candidate standard, the 95-percent uncertainty for the corrected concentration must be included in the assessment of the total analytical uncertainty of the candidate standard's concentration using the following equation:

$$Uncertainty_{TOTAL} = \sqrt{(Uncertainty_{ASSAY})^2 + (Uncertainty_{CORRECTION})^2 + (Uncertainty_{STANDARD})^2} .$$

The analyst may report the total analytical uncertainty of the candidate standard's certified concentration on the certification documentation or may report this estimate as a percentage that is relative to the certified concentration using the following equation:

$$Uncertainty_{RELATIVE} = 100 \left[Uncertainty_{TOTAL} / Certified Concentration \right].$$

2.1.9 Zero Gas

Zero gas used for zero gas checks or for dilution of any candidate or reference standard should be clean, dry, zero-grade air or nitrogen containing no detectable concentration of the pollutant of interest. It should match the balance gas in the candidate standard and the reference standard, unless it has been demonstrated that the analyzer is insensitive to differences in the balance gas composition. The zero gas also should contain no contaminant that causes a detectable response on the analyzer or that suppresses or enhances the analyzer's response. The oxygen content of zero air should be approximately that of ambient air, unless it has been demonstrated that varying the oxygen content does not suppress or enhance the analyzer's response. The water vapor concentration in the zero gas should be less than 5 ppm.

The analyst may substitute a low-concentration, NIST-traceable reference standard for the zero gas in zero gas checks and assays if there is reason to believe that the zero gas reading may not accurately represent the zero-intercept of the calibration equation.

2.1.10 Accuracy Assessment of Commercially Available Standards

Periodically, the U.S. EPA will assess the accuracy of compressed gas calibration standards that have been assayed and certified according to this protocol. The accuracy of representative standards will be assessed by EPA audits. The audit results, identifying the specialty gas producers or other analytical laboratories that assayed and certified the standards, will be published as public information. A summary of EPA's audit results from 1992 to the present is available as a WordPerfect 6.1 file at EPA's Ambient Monitoring Technology Information Center (Internet Address: http://www.epa.gov/ttn). This document can be found in the Directory of TTNWeb Sites/ AMTIC/Publications/ORD-NERL Documents.

2.2 PROCEDURE G1: ASSAY AND CERTIFICATION OF A COMPRESSED GAS CALIBRATION STANDARD WITHOUT DILUTION

2.2.1 Applicability

This procedure may be used to assay the concentration of a candidate compressed gas calibration standard, based on the concentration of a compressed gas reference standard of the same gas mixture. This procedure allows a specialty gas producer, a standard user, or other analytical laboratory to certify that the assayed concentration for the candidate standard is traceable to the reference standard. The procedure employs a pollutant gas analyzer to compare the candidate and reference standards' concentrations by direct measurement without dilution of either gas.

This procedure may be used for the assay of more than one candidate standard during the same assay session. Criteria that apply to the assay of one candidate standard apply to the assay of multiple candidate standards.

2.2.2 Limitations

The concentration of the candidate standard may be greater than or lesser than the concentration of the reference standard. However, both concentrations must lie within the well-characterized region of the multipoint calibration (see Subsection 2.1.7.2). Additionally, the magnitude of the 95-percent confidence limits for the estimated concentration of the candidate standard must $\leq \pm 1$ percent of the reference standard concentration. This criterion may be more restrictive than the corresponding criterion for the multipoint calibration, but it allows the analyst greater flexibility in the selection of a reference standard for the assay of a particular candidate standard. For example, assume that a 70-ppm candidate standard is being assayed using a 50-ppm reference standard and that the analyzer's calibration was found to be well-characterized between 20 and 80 ppm. The 95-percent confidence limits for the candidate standard's estimated concentration must be less than or equal to ± 0.5 ppm.

The balance gas must be the same in both the candidate standard and the reference standard, unless it has been demonstrated that the analyzer's response is insensitive to differences in the balance gas composition.

2.2.3 Assay Apparatus

Figure 2-2 illustrates one possible design of apparatus for the assay of compressed gas calibration standards without dilution. This apparatus is designed to allow the convenient routing of the gas mixtures to the pollutant gas analyzer. The gas mixture to be measured is selected by rotation of two three-way valves (i.e., V1 and V2). Pressure regulators and gas flow controllers (i.e., C1 and C2) control the flow rates from the individual cylinders. The gas flow controllers may be needle values, capillary tubes, thermal mass flow controllers, or other flow control devices. The gas mixtures are routed to the analyzer through a union tee tube fitting. Gas in excess of the analyzer's demand is vented, which helps to ensure that the gas entering the analyzer is at near-ambient pressure. Normally, the excess gas is vented to the atmosphere without any obstructions in the tubing. However, the excess gas can be routed through an uncalibrated rotameter by rotation of a three-way valve (i.e., V3). The rotameter is used to demonstrate that the total gas flow rate exceeds the sample flow rate of the analyzer and that no room air is being drawn in through the vent line.

The apparatus may be modified in several ways that will not diminish its performance. The two three-way valves could be replaced by solenoid valves or by a single four-way valve with three input ports and one output port. Alternatively, a single length of tubing with a gas flow controller could be connected manually to individual cylinders' pressure regulators in succession. See also Subsection 2.1.3.

2.2.4 Pollutant Gas Analyzer

The pollutant gas analyzer must have a well-characterized calibration curve and must be capable of measuring directly the concentration of both the candidate and the reference standards without dilution. See Subsection 2.1.7.1. It must have good resolution, good precision, a stable response, and low output signal noise. In addition, the analyzer should have good specificity for the pollutant of interest so that it has no detectable response to any other component or contaminant that may be contained in either the candidate or reference standards. If any component in a multiple-component standard interferes with the assay of any other component, the analyst must conduct an interference study to determine an interference correction equation. If the candidate and reference standards contain dissimilar balance gases (e.g., air versus nitrogen or different pro-portions of oxygen in the balance air), it must have been demonstrated that the analyzer's response is not sensitive to difference exists in the balance gas composition. This demonstration can be accomplished by showing that no difference exists in the analyzer's response when measuring a compressed gas calibration standard that has been diluted with identical flow rates of different balance gases.

The analyzer should be connected to a high-precision data acquisition system (e.g., a strip chart recorder), which must produce an electronic or paper record of the analyzer's response during the assay. A high-precision digital panel meter, a digital voltmeter, a data logger or some other data acquisition system with four-digit resolution can be used to obtain numerical values of the analyzer's response. More precise values will be obtained if this system has a data-averaging capability. The assay record must be maintained for 3 years after the standard's certification date.

If the analyzer has not been in continuous operation, turn it on and allow it to stabilize (e.g., for at least 12 hours) before beginning the measurements.

2.2.5 Analyzer Calibration

2.2.5.1 Multipoint Calibration—

See Subsections 2.1.7.2 and 2.1.7.4.

2.2.5.2 Analyzer Range—

The range of the analyzer must include the concentrations of the zero gas, the candidate standard and the reference standard. The concentrations of the candidate and reference standards must fall within the well-characterized region of the analyzer's calibration curve. In general, the analyst should use a range that will produce the largest on-scale analyzer response.

2.2.5.3 Linearity—

The data reduction technique used in this procedure requires that the analyzer have a well-characterized, but not necessarily linear, calibration curve (see Subsection 2.1.7.5). High-concentration-range analyzers of the type that are required for this procedure may not be inherently linear, but in such cases they usually have a predictable, non-linear calibration curve that can be described by a polynomial equation or can be mathematically transformed to produce a straight-line calibration curve that is suitable for use in this procedure. Any such polynomial equation or mathematical transformation should be verified during the multipoint calibration. Caution should be

exercised in using a transformed calibration curve because zero or span control adjustments to the analyzer may produce unexpected effects in the transformed calibration curve.

2.2.5.4 Zero and Span Gas Checks—

See Subsections 2.1.7.3 and 2.1.7.4. Prior to carrying out the assay of the candidate standard, use zero and span gases to check for calibration drift in the analyzer since the multipoint calibration. Zero gas and span gas checks must be performed on any day after the multipoint calibration that candidate standards are assayed. If multiple assays are being performed on the same analyzer range, the analyst needs to perform only a single set of zero gas and span gas checks for this range. However, another set must be performed if the range is changed.

The gas mixtures to be used during the zero and span gas checks need not be the same as any of the reference standards used for the assay of the candidate standard or for the multipoint calibration. The reference standard for the span gas check must be traceable to a NIST SRM, a SRM-equivalent PRM, or an NTRM. Information concerning this standard (e.g., cylinder identification number, certified concentration) must be recorded in the laboratory's records.

A source of clean, dry zero gas is recommended, but not required. The analyst may substitute a low-concentration, NIST-traceable reference standard for the zero gas if there is reason to believe that the zero gas reading may not accurately represent the zero-intercept of the calibration equation.

Make three or more discrete measurements of the zero gas and three or more discrete measurements of the reference standard. "Discrete" means that the analyst must change the gas mixture being sampled by the analyzer between measurements. For example, the analyst might alternate between measurements of the reference standard and measurements of the zero gas. Record these measurements in the laboratory's records.

Next, verify that the analyzer's precision is acceptable. Calculate the mean and standard deviation of the analyzer's responses to the zero gas. Repeat the calculations for the reference standard measurements. These calculations are performed in Step 6 of the spreadsheet described in Appendix A. Record these calculations in the laboratory's records. The standard error of the mean for each set of measurements must be less than or equal to 1.0 percent of the mean response to the reference standard. That is,

$$\frac{s}{\sqrt{n}} \le \frac{\overline{R_{RS}}}{100}$$

where

s = standard deviation of the analyzer's response;

n = the number of independent measurements of the gas mixture; and

 $\overline{R_{RS}}$ = the mean analyzer response to the reference standard.

The value of the standard error of the mean can be made smaller by increasing the number of measurements. This calculation will enable the analyst to determine how many replicate measurements are needed during the assay of the candidate standard to obtain acceptable precision. The analyst may wish to use a data logger or data acquisition system with averaging capability to obtain more precise measurements. If the value of the standard error of the mean is not acceptable, then the analyzer must be repaired or another analyzer must be used for the assay.

Next, verify that excessive calibration drift has not occurred since the multipoint calibration. For the zero gas measurements, calculate the relative difference (in percent) between the current mean

analyzer response during the zero gas check and the corresponding response that is predicted from the multipoint calibration regression equation. That is,

Relative Difference =
$$100 \left[\frac{\text{Current Response} - \text{Calibration Response}}{\text{Calibration Response for Reference Standard}} \right]$$
.

This calculation is performed in Step 6 of the spreadsheet described in Appendix A.

Note that the relative difference is always calculated relative to the calibration response for the reference standard, even when the zero gas is being measured. Repeat this calculation for the reference standard measurements. Record these calculations in the laboratory's records. If the reference standard was not measured during the multipoint calibration, use the regression-predicted response for a concentration equal to that of the reference standard.

Then, if the relative differences for the zero and span gas checks are less than or equal to 5.0 percent, the analyzer is considered to be sufficiently in calibration. The zero and span controls need not be adjusted and the assay may be conducted. The data reduction technique does not require the absolute accuracy of the analyzer calibration. Some minor calibration drift is acceptable because the effect of any drift will be corrected during the reduction of the assay data.

However, if the relative difference for either the zero or the span gas check is greater than 5.0 percent, then the analyzer is considered to be out of calibration. A new multipoint calibration may be conducted before the candidate standard is assayed or the analyzer's zero and span controls may be adjusted to return the analyzer's response to the original calibration levels. For some analyzers such as nondispersive infrared instruments, daily changes in environmental variables such as barometric pressure may shift the calibration. After any adjustment of controls, the analyst should repeat the zero and span gas checks and recalculate the relative differences to verify that the analyzer is sufficiently in calibration. The analyzer will be considered to be out of calibration if the relative differences remain greater than 5.0 percent.

The zero gas and reference standard measurements that are performed for the assay of the candidate standard may also be used for the zero and span gas checks.

2.2.6 Assay Gases

2.2.6.1 Candidate Standard—

See Subsections 2.1.6 and 2.2.2.

2.2.6.2 Reference Standard—

See Subsections 2.1.2 and 2.2.2. The reference standard used for the assay of the candidate standard must be a NIST SRM, an SRM-equivalent PRM, an NTRM or a GMIS. This standard need not be the same as any of the reference standards used for the span gas check or for the multipoint calibration. Information concerning the reference standard (e.g., cylinder identification number, certified concentration, etc.) must be recorded in the laboratory's records.

If the multipoint calibration data have been fitted to a linear (i.e., straight-line) regression model, then only a single reference standard need be measured during the assay of the candidate standard. If these data have been fitted to a quadratic or higher-order regression model, then at least two reference standards must be measured. One reference standard is adequate to determine the slope of a linear equation, but additional reference standards are needed to determine the curvature of quadratic and higher-order polynomial equations. The concentrations of the additional reference standards should be

located at the maximum difference between the polynomial curve and the corresponding straight line between the zero gas and the highest-concentration reference standard.

2.2.6.3 Zero Gas—

See Subsection 2.1.9. A source of clean, dry zero gas is recommended, but not required. The analyst may substitute a low-concentration, NIST-traceable reference standard for the zero gas during zero gas checks and assays if there is reason to believe that the zero gas reading may not accurately represent the zero-intercept of the calibration equation. Information concerning the zero gas should be recorded in the laboratory's records.

2.2.7 Assay Procedure

- Verify that the assay apparatus is properly configured, as described in Subsection 2.2.3 and shown in Figure 2-1. Inspect the analyzer to verify that it appears to be operating normally and that all controls are set to their expected values. Record these control values in the laboratory's records.
- 2. Verify that a multipoint calibration of the analyzer has been performed within 1 month prior to the assay date. (See Subsections 2.1.7.2, 2.1.7.5 and 2.2.4). Additionally, verify that the zero and span gas checks indicate that the analyzer is in calibration (see Subsection 2.2.5.4). Finally, verify that the concentrations of the candidate and reference standards fall within the well-characterized region of the analyzer's calibration curve (see Subsection 2.2.2).
- 3. Measure and adjust the flow rates of the gas mixtures (i.e., reference standard(s), candidate standard, and zero gas) to approximately the same value that will provide enough flow for the analyzer and sufficient excess to assure that no ambient air will be drawn into the vent line.
- 4. In succession, measure the zero gas, the reference standard(s), and the candidate standard(s) using the analyzer. Use valves V1 and V2 to select each of the gas mixtures for measurement. For each measurement, allow ample time for the analyzer to achieve a stable response. If the response for each measurement is not stable, the precision of the measurements will decline and the candidate standard may not be certifiable under this protocol. Record the analyzer response for each measurement in the laboratory's records, using the same response units (e.g., volts, millivolts, percent of scale, etc.) as was used for the multipoint calibration. At this point, do not convert these data into concentration values using the calibration equation. Do not perform any necessary mathematical transformation of these data. These steps will be done later. Do not make any zero control, span control, or other adjustments to the analyzer during these measurements.

The analyst may assay multiple candidate standards during the same assay session. For example, a single set of measurements may involve a zero gas, a reference standard and three candidate standards. Criteria that apply to the assay of one candidate standard apply to the assay of multiple candidate standards. The analyst should be aware that the effect of any short-term calibration drift will be greater when multiple candidate standards are assayed. This greater effect is due to the longer period of time between reference standard measurements. Unacceptable uncertainties of the estimated concentrations for the candidate standards may occur as a result of the longer assay session.

5. Conduct at least two additional sets of measurements, as described in step 4 above. However, for these subsequent sets of measurements, change the order of the three measurements (e.g., measure reference standard, zero gas, and candidate standard for the second set and measure zero gas, candidate standard, and reference standard for the third set). Changing the order

that the gas mixtures are measured helps the analyst to discover any effect of that one measurementhas on subsequent measurements. The number of sets of measurements will have been determined during analysis of the multipoint calibration data such that the 95-percent uncertainty for the regression-predicted concentration of the candidate standard is ≤ 1 percent of the concentration of the reference standard.

- 6. If any one or more of the measurements of a set of measurements is invalid or abnormal for any reason, discardall three measurements and repeat the measurements. Such measurements may be discarded if the analyst can demonstrate that the experimental conditions were inappropriate during these measurements. Data cannot be discarded just because they appear to be outliers but may be discarded if they satisfy statistical criteria for testing outliers. The analyst must record any discarded data and a brief explanation as to why these data were discarded in the laboratory's records.
- 7. The spreadsheet described in Appendix A or equivalent statistical techniques must be used to calculate an estimated concentration and a 95-percent uncertainty for the candidate standard based on data from the assay measurements and from the multipoint calibration. The use of both sets of data in the statistical analysis produces an estimated concentration with smaller uncertainty while correcting for any minor calibration drift which may have occurred since the multipoint calibration. Record the estimated concentration and the 95-percent uncertainty in the laboratory's records.

The spreadsheet also calculates the percentage of the uncertainty that is due to the multipoint calibration. This percentage is needed for the total uncertainty calculations when two or more assays fall under the same multipoint calibration. Record this value in the laboratory's records.

The analyst should investigate any of the measurements that appear to be outliers. Such data may be discarded if the analyst can demonstrate that the experimental conditions were inappropriate during these measurements. Data cannot be discarded just because they appear to be outliers, but may be discarded if they satisfy statistical criteria for testing outliers. The analyst must record any discarded data and a brief summary of the investigation in the laboratory's records.

8. If the multipoint calibration data and the assay data underwent any mathematical transformations before their statistical analysis, perform the reverse transformations for the estimated concentration and the 95-percent uncertainty. Record the transformed values in the laboratory's records.

2.2.8 Stability Test for Newly Prepared Candidate Standards

Newly prepared candidate standards that contain reactive gas mixtures must be assayed on at least two dates that are separated by at least 7 days (see Subsections 2.1.6.1 and 2.1.6.2). The spreadsheet described in Appendix C or equivalent statistical techniques must be used to evaluate the stability of the candidate standard and to calculate the overall estimated concentration and the total uncertainty for the candidate standard.

The stability is evaluated by comparing the 95-percent confidence limits (i.e., estimated concentration ±95-percent uncertainty) for the candidate standard from the two or more assays. If the confidence limits overlap, the candidate standard can be considered to be stable and may be certified. In the spreadsheet, all cells in the comparisons table will be "true." If the confidence limits do not overlap, the candidate standard may be unstable or there may be analytical problems associated with the assays or the reference standards. One or more cells in the comparisons table will be "false." The

analyst must either disqualify the candidate standard or investigate why the confidence limits do not overlap. The analyst may conduct additional assays until stability is achieved and add the additional data to the spreadsheet. Data from a nonoverlapping assay may be discarded and the remaining data used to determine the overall estimated concentration and the total uncertainty provided the confidence limits overlap. Record these values and any discarded data in the laboratory's records.

2.2.9 Certification Documentation

See Subsections 2.1.4 and 2.1.5.

2.2.10 Recertification Requirements

See Subsections 2.1.6.3 and 2.1.6.4.

2.3 PROCEDURE G2: ASSAY AND CERTIFICATION OF A COMPRESSED GAS CALIBRATION STANDARD USING DILUTION

2.3.1 Applicability

This procedure may be used to assay the concentration of a diluted candidate compressed gas calibration standard, based on the concentration of a diluted compressed gas reference standard of the same gas mixture. This procedure allows a specialty gas producer, a standard user, or other analytical laboratory to certify that the assayed concentration for the candidate standard is traceable to the reference standard. The procedure employs a low-concentration-range (i.e., ambient air quality level) pollutant gas analyzer to compare quantitatively diluted gas samples of both the candidate and reference standards.

Dilution of the candidate and reference standards with zero gas allows greater flexibility in the range of concentrations of both the candidate and reference standards that can be assayed. Additionally, dilution allows the use of a low-concentration-range analyzer, which is more likely to have an inherently linear response than a high-concentration-range analyzer. However, the dilution technique introduces additional error into the assay which would not be present if the standards were assayed without dilution. This additional error is measured by an accuracy check of the assay apparatus which is performed as part of the multipoint calibration.

This procedure may be used for the assay of multiple candidate standards at the same time. Criteria that apply to the assay of one candidate standard apply to the assay of multiple candidate standards.

2.3.2 Limitations

The concentration of the diluted candidate standard may be greater than or lesser than the
concentration of the diluted reference standard. However, both concentrations must lie
within the well-characterized region of the analyzer's multipoint calibration (see
Subsection 2.1.7.2).

Additionally, the magnitude of the 95-percent confidence limits for the estimated concentration of the candidate standard must be ≤±1 percent of the reference standard concentration. This criterion may be more restrictive than the corresponding criterion for the multipoint calibration, but it allows the analyst greater flexibility in the selection of a reference standard for the assay of a particular candidate standard. For example, assume that a 70-ppm candidate standard is being assayed using a 50-ppm reference standard and that the analyzer's calibration was found to be well characterized between 20 and 80 ppm. The 95-

percent confidence limits for the candidate standard's estimated concentration must be $\leq \pm 0.5$ ppm.

- 2. An accurate system for flow measurement and gas dilution is required.
- 3. The balance gas in both the candidate and reference standards must be identical, unless either a high dilution flow rate ratio (i.e., at least 50 parts zero gas to 1 part standard) is used for the assay or it has been demonstrated that the analyzer is insensitive to differences in the balance gas.

2.3.3 Assay Apparatus

The components of the assay apparatus can be assembled in several different configurations without diminishing performance. Two possible designs of the assay apparatus are illustrated in Figures 2-3 and 2-4. The former figure shows a configuration in which discrete components (i.e., three-way valves, gas flow controllers, and a mixing chamber) are used to dilute the reference and candidate standards. The latter figure shows a configuration in which a commercially available gas dilution system is used to dilute the standards. Both designs share the important characteristic that the candidate standard is diluted by the same components that dilute the reference standard.

In Figure 2-3, either zero gas or a diluted standard can be routed to the analyzer by rotation of three three-way values (i.e., V1, V2, and V3). One gas flow controller (i.e., C1) regulates the flow rates of the reference and candidate standards. These flow rates can be measured by a single flowmeter connected to an outlet port on valve V2 or by a flowmeter built into C1. Another gas flow controller (i.e., C2) regulates the flow rate of the zero gas. This flow rate can be measured by a flowmeter connected to an outlet port on valve V3 or by a flowmeter built into C2. The gas flow controllers may be needle valves, capillary tubes, thermal mass flow controllers, or other suitable devices (see Subsection 2.3.7). If different flow rates are used for the reference and candidate standards during the assay (see Subsection 2.3.6), separate gas flow controllers may be used for the two standards. However, the same flowmeter must be used to measure the two flow rates to minimize error in the measurement (see Subsection 2.3.7). Flow rates should be controlled and measured with a relative uncertainty of 1 percent or less. For large dilutions of the standards, the reference and candidate standard flow rates may be quite small. Therefore, the internal volume of the tubing and components should be kept small to minimize the flushing time when valve V1 is rotated.

The mixing chamber combines the two gas streams and should be designed to produce turbulent flow to ensure thorough mixing of the gas streams. The diluted gas mixtures are routed to the analyzer through a union tee tube fitting, which vents excess gas flow. Normally, the excess gas is vented to the atmosphere without any obstructions in the tubing and the gas entering the analyzer is at near-atmospheric pressure. However, the excess gas can be routed through an uncalibrated rotameter by rotation of a three-way valve (i.e., V4). The rotameter is used to demonstrate that the total gas flow rate exceeds the sample flow rate of the analyzer and that no room air is being drawn in through the vent line.

The apparatus in Figure 2-3 may be modified in several ways that will not diminish its performance. For example, the three-way valves could be replaced by solenoid valves. Alternatively, valve V1 could be replaced by a single length of tubing that is connected manually to the two standards' pressure regulators in succession (see also Subsection 2.1.3).

In Figure 2-4, the reference and candidate standards are diluted with a gas dilution system. This gas dilution system may use capillary tubes, positive-displacement pumps, thermal mass flow controllers, or other suitable devices to dilute the standards. If a gas dilution system is used, it must have a specified accuracy of not greater than 1.0 percent of the undiluted reference standard concentration. The analyst must check the accuracy of the gas dilution system during the multipoint calibration (see Subsections 2.1.7.2 and 2.3.5.1).

2.3.4 Pollutant Gas Analyzer

The pollutant gas analyzer must have a well-characterized calibration curve and must have a range that is capable of measuring the diluted concentration of both the candidate and the reference standards (see Subsection 2.1.7.1). It must have good resolution, good precision, a stable response, and low output signal noise. In addition, the analyzer should have good specificity for the pollutant of interest so that it has no detectable response to any other component or contaminant that may be contained in either the candidate or reference standards. If any com-ponent in a multiple-component standard interferes with the assay of any other component, the analyst must conduct an interference study to determine an interference correction equation. A suitable analyzer with acceptable performance specifications may be selected from the list of EPA-designated reference and equivalent method analyzers.¹⁷ If the candidate and reference standards contain dissimilar balance gases (e.g., air versus nitrogen or different proportions of oxygen in the balance air), either a high dilution flow-rate ratio (i.e., at least 50 parts zero gas to 1 part standard) should be used or it must have been demonstrated that the analyzer's response is not sensitive to differences in the balance gas composition. This demonstration may be accomplished by showing that no difference exists in an analyzer's response when measuring a compressed gas calibration standard that has been diluted with identical flow rates of different balance gases.

The analyzer should be connected to a high-precision data acquisition system (e.g., a strip chart recorder) which must produce an electronic or paper record of the analyzer's response during the assay. A high-precision digital panel meter, a digital voltmeter, a data logger or some other data acquisition system with four-digit resolution can be used to obtain numerical values of the analyzer's response. More precise values will be obtained if this system has a data-averaging capability. The assay record must be maintained for 3 years after the standard's certification date.

If the analyzer has not been in continuous operation, turn it on and allow it to stabilize (e.g., for at least 12 hours) before beginning any measurements.

2.3.5. Analyzer Calibration

2.3.5.1 Multipoint Calibration—

See Subsections 2.1.7.2 and 2.1.7.4. Following completion of the multipoint calibration, the accuracy of the assay apparatus must be checked to verify that the error associated with the dilution is not excessive. This accuracy check involves the measurement of an undiluted or diluted check standard. The check standard must be an SRM, an SRM-equivalent PRM, or an NTRM, or a GMIS as specified in Subsection 2.1.2. It must have a certified concentration that is different from that of the reference standard used in the multipoint calibration. Information concerning this standard (e.g., cylinder identification number, certified concentration) must be recorded in the laboratory's records.

If an undiluted check standard is used, its concentration must fall in the well-characterized region of the calibration curve. If a diluted check standard is used, the diluted concentration must fall in the well-characterized region.

Make three or more discrete measurements of the undiluted or diluted check standard. "Discrete" means that the analyst must change the gas mixture being sampled by the analyzer between measurements. For example, the analyst might alternate between measurements of the check standard and the zero gas. Record these measurements in the laboratory's records.

Next the analyst must verify that the dilution error is not excessive. For the check standard measurements, calculate the relative difference (in percent) between the mean analyzer response and the corresponding response that is predicted from the multipoint calibration regression equation and the undiluted or diluted check standard concentration. That is.

Relative Difference =
$$100 \left[\frac{\text{Mean Analyzer Response} - \text{Predicted Response}}{\text{Predicted Response}} \right]$$
.

If the relative difference is greater than 1.0 percent, the dilution error is considered to be excessive. The analyst must investigate why the relative difference is excessive. The problem may be due to errors in the reference standard and check standard concentrations, errors in assay apparatus or to some other source. Assays may not be conducted until the relative difference for a subsequent accuracy check is less than or equal to 1.0 percent.

2.3.5.2 Analyzer Range—

The range of the analyzer must include the concentrations of the zero gas and of the diluted candidate and reference standards (see Subsection 2.3.6). The concentrations of the diluted reference and candidate standards must fall within the well-characterized region of the analyzer's calibration curve. Because the selection of the dilution ratio or ratios to be used in the assay provides great flexibility in the choice of concentrations to be measured by the analyzer, the analyzer range should be selected based on optimum accuracy, stability, and linearity.

2.3.5.3 Linearity—

The data reduction technique used in this procedure requires that the analyzer have a well-characterized, but not necessarily linear, calibration curve (see Subsection 2.1.7.5). Many lower-concentration analyzers of the type that may be used for this procedure have straight-line calibration curves. If not, they usually have a predictable nonlinear calibration curve that can be described by a polynomial equation or can be mathematically transformed to produce a straight-line calibration curve suitable for use in this procedure. Any such polynomial equation or mathematical transformation should be verified during the multipoint calibration. Caution should be exercised in using a transformed calibration curve because zero or span control adjustments to the analyzer may produce unexpected effects in the transformed calibration curve.

2.3.5.4 Zero and Span Gas Checks—

See Subsections 2.1.7.3 and 2.1.7.4. Prior to carrying out the assay of the candidate standard, use zero and span gases to check for calibration drift in the analyzer since the multipoint calibration. Zero gas and span gas checks must be performed on any day after the multipoint calibration that candidate standards are assayed. If multiple assays are being performed on the same analyzer range, the analyst needs to perform only a single set of zero gas and span gas checks. However, another set must be performed if the range is changed.

The gas mixtures to be used during the zero and span gas checks need not be the same as any of the reference standards used for the assay of the diluted candidate standard or for the multipoint

calibration. The reference standard for the span gas check must be traceable to a NIST SRM, an SRM-equivalent PRM, or an NTRM. Information concerning this standard (e.g., cylinder identification number, certified concentration) must be recorded in the laboratory's records.

Make three or more discrete measurements of the zero gas and three or more independent measurements of the diluted reference standard. Record these measurements in the laboratory's records.

Next, the analyst must verify that the analyzer's precision is acceptable. Calculate the mean and standard deviation of the analyzer's response to the zero gas. Repeat these calculations for the diluted reference standard measurements. These calculations are performed in Step 6 of the spreadsheet described in Appendix A. Record these calculations in the laboratory's records. The standard error of the mean for each set of measurements must be less than or equal to 1.0 percent of the mean response to the diluted reference standard. That is,

$$\frac{s}{\sqrt{n}} \le \frac{\overline{R}_{DRS}}{100}$$

where

s = standard deviation of the analyzer's response;

n = the number of independent measurements of the gas mixture; and

 $\frac{1}{R_{DDS}}$ = the mean analyzer response to the diluted reference standard.

The value of the standard error of the mean can be made smaller by increasing the number of measurements. This calculation will enable the analyst to determine how many replicate measurements are needed during the assay of the diluted candidate standard to obtain acceptable precision. The analyst may wish to use a data logger or data acquisition system with data averaging capability to obtain more precise measurements. If the value of the standard error of the mean is not acceptable, then the analyzer must be repaired or another analyzer must be used for the assay.

Next the analyst must verify that excessive calibration drift has not occurred since the multipoint calibration. For the zero gas measurements, calculate the relative difference (in percent) between the mean analyzer response during the zero gas check and the corresponding response that is predicted from the multipoint calibration regression equation. That is,

Relative Difference =
$$100 \left[\frac{\text{Current Response} - \text{Calibration Response}}{\text{Calibration Response for Diluted Reference Standard}} \right]$$
.

This calculation is performed in Step 6 of the spreadsheet described in Appendix A.

Note that the relative difference is always calculated relative to the calibration response for the diluted reference standard, even when zero gas is being measured. Repeat this calculation for the diluted reference standard measurements. Record these calculations in the laboratory's records. If the diluted reference standard was not measured during the multipoint calibration, use the regression-predicted response for a concentration equal to that of the diluted reference standard.

Then, if the relative differences for the zero and span checks are less than or equal to 5.0 percent, the analyzer is considered to be in calibration. The zero and span controls need not be adjusted and the assay may be conducted. The data reduction technique used in this procedure does not require the

absolute accuracy of the analyzer's calibration. Some minor calibration drift is acceptable because the drift will be corrected for during the reduction of the assay data.

However, if the relative difference for either the zero or the span gas checks is greater than 5.0 percent, then the analyzer is considered to be out of calibration. A new multipoint calibration may be conducted before the candidate standard is assayed or the analyzer's zero and span controls may be adjusted to return the analyzer's response to the original calibration levels. For some analyzers such as nondispersive infrared instruments, daily changes in environmental variables such as barometric pressure may shift the calibration. After any adjustment of the controls, the analyst should repeat the zero and span gas checks and recalculate the relative differences to verify that the analyzer is sufficiently in calibration. The analyzer will be considered to be out of calibration if the relative differences remain greater than 5.0 percent.

The zero gas and diluted reference standard measurements that are performed for the assay of the diluted candidate standard may also be used for the zero gas and span gas checks.

2.3.6 Selection of Gas Dilution Flow Rates or Gas Concentration Settings

The flow rates or settings used for the zero gas, reference standard, and candidate standard should be selected carefully to provide diluted concentrations for both the candidate and reference standards that fall in the well-characterized region of the analyzer's calibration curve. The diluted concentration of the candidate standard may be greater than or lesser than the diluted concentration of the reference standard. Any assay error due to the dilution process will be reduced if the same dilution flow-rate ratio or concentration setting can be used for both the candidate and reference standards. Select the diluted concentrations of the reference and candidate standards, and select flow rates or concentration settings that will produce the highest analyzer responses within the well-characterized region of the analyzer's calibration curve.

If the same dilution flow-rate ratio or concentration setting cannot be used for both the candidate and reference standards, select different ratios or settings for the candidate and reference standards to produce concentrations that are approximately equal and that fall in the well-characterized region of the analyzer's calibration curve. Select flow rates or settings such that only one of the apparatus controls must be adjusted when switching from the reference standard to the candidate standard, or vice versa. Where a choice of analyzer ranges is available, higher dilution ratios or lower concentration settings will reduce the consumption of the standards.

2.3.7 Flowmeter Type and Flowmeter Calibration

Figure 2-3 shows flow measurement ports on valves V2 and V3. In this configuration, a single flowmeter can be used to measure both the standard flow rate and the zero gas flow rate. Such an approach would reduce measurement errors arising from differences in the calibration of multiple flowmeters. Alternatively, the flow rates can be measured at the outlet of the dilution apparatus, with the rotameter vent temporarily plugged. In either case, a NIST-traceable volumetric flowmeter such as a wet test meter, a thermal mass flowmeter, or a soap bubble flowmeter can be used (see Subsection 2.1.2). Each flow rate must be measured separately while the other flow rates are set to zero. Care must be exercised to ensure that each measured flow rate remains constant when combined with the other flow rate(s) and between the time of measurement and the time of the assay. Additionally, care must be taken to ensure that the flowmeter does not cause any back pressure in the gas stream and any resulting change in the flow rate through the flow controller.

If in-line flowmeters are mounted directly downstream of the flow controllers, they may not operate at atmospheric pressure because of back pressure from downstream components. Also, this back pressure may vary as a function of the total flow rate. Thus, the flowmeters must compensate for the variable in-line pressure. Thermal mass flowmeters do not need to be corrected for pressure effects.

as wet test meters must be carefully corrected for the actual gas pressure during the flow measurement. An in-line flowmeter must not contaminate or react with the gas mixture passing through it.

The flowmeters used should be stable, repeatable, and linear and have good resolution. If possible, select flow rates or a flowmeter range such that the flow rates to be measured fall in the upper half of the flowmeter's range. The flowmeters should be carefully calibrated at several flow rates to prove linearity. The calibration should be accurate to plus or minus 1 percent and must be referenced to an accurate flow rate or volumetric standard traceable to a NIST primary standard. Flowmeter calibrations should be checked and recertified periodically, as determined by stability information such as a chronological control chart of calibration data.

All volumetric flow-rate measurements must be corrected or referenced to the same temperature and pressure conditions, such as EPA-standard conditions (i.e., 760 millimeters of mercury (mm Hg), 25 °C) or the ambient temperature and pressure conditions prevailing in the laboratory during the assay. Measurements using wet test meters and soap bubble flowmeters also must be corrected for the saturation of the gas stream with water vapor in the moist interiors of these flowmeters. The equation to correct the flow rate for temperature, pressure, and humidity effects is given below:

Flow Rate =
$$\frac{\text{Volume}}{\text{Time}} \left(\frac{P_{\text{M}} - P_{\text{WV}}}{P_{\text{S}}} \right) \left(\frac{T_{\text{S}}}{T_{\text{M}}} \right)$$

where

measured barometric pressure (mm Hg); partial pressure of water vapor (mm Hg);

standard pressure (mm Hg);

standard temperature (298.2 K); and

measured ambient temperature (273.2 + °C).

Measurement of reference and candidate standard flow rates with the same flowmeter and measurement of both dilution zero gas flow rates with the same flowmeter tend to reduce measurement errors, associated with the use of multiple flowmeters. These errors are more pronounced at higher dilution flow rate ratios. Note that the impact of any flow measurement error is reduced if the same dilution ratio can be used for both the reference standard and candidate standard measurements.

2.3.8 **Assay Gases**

2.3.8.1 Candidate Standard—

See Subsections 2.1.6, 2.3.2, and 2.3.6.

2.3.8.2 Reference Standard—

See Subsections 2.1.2, 2.1.6.4, 2.3.2, and 2.3.6. The reference standard used for the assay of the candidate standard must be a NIST SRM, an SRM-equivalent PRM, an NTRM or a GMIS. This standard need not be the same as any of the reference standards used for the span gas check or for the multipoint calibration. Information concerning the reference standard (e.g., cylinder identification number, certified concentration, etc.) must be recorded in the laboratory's records.

If the multipoint calibration data have been fitted to a linear (i.e., straight-line) model, then only a single reference standard need be measured during the assay of the candidate standard. If these data have been fitted to a quadratic or higher-order polynomial model, then at least two reference standards must be measured. One reference standard is adequate to determine the slope of a linear equation, but additional reference standards are needed to determine the curvature of quadratic or higher-order polynomial equations. The concentrations of the additional reference standards should be located at the maximum difference between the polynomial curve and the corresponding straight line between the zero gas and the highest-concentration reference standard.

2.3.8.3 Zero Gas—

See Subsection 2.1.9. Use the same zero gas for dilution of both candidate and reference gases. The analyst may substitute a low-concentration, NIST-traceable reference standard for the zero gas in zero gas checks and assays if there is reason to believe that the zero gas reading may not accurately represent the zero-intercept of the calibration equation. Information concerning the zero gas should be recorded in the laboratory's records.

2.3.9 Assay Procedure

- Verify that the assay apparatus is properly configured as shown in Figure 2-3 or Figure 2-4 and as described in Subsection 2.3.3. Inspect the analyzer to verify that it appears to be operating normally and that all controls are set to their expected values. Record these control values in the laboratory's records.
- 2. Verify that the flowmeters, if used in the assay apparatus, are properly calibrated (see Subsection 2.3.7).
- 3. Verify that a multipoint calibration of the analyzer has been performed within 1 month prior to the assay date and that the dilution error is not excessive (see Subsections 2.1.7.2, 2.1.7.5, 2.3.4, and 2.3.5.1). Additionally, verify that the zero and span gas checks indicate that the analyzer is in calibration (see Subsection 2.3.5.4). Finally, verify that the concentrations of the diluted reference and candidate standards fall within the well-characterized region of the analyzer's calibration curve (see Subsection 2.3.2).
- 4. Determine and establish the flow rates or concentration settings of the gas mixtures (i.e., reference standard(s), candidate standard, and zero gas) that will be used for the assay (see Subsections 2.3.6, 2.3.7, and 2.3.5.2). Also check that the total flow rate coming from the mixing chamber will provide enough flow for the analyzer and sufficient excess to ensure that no ambient air will be drawn into the vent line. Changes in the sample pressure may change the calibration curve. When using the same flow rates for both candidate and reference standards, carefully set the delivery pressures of the two standards' pressure regulators to the same value so that there is no change in the flow rate when switching from one standard to the other.

Calculate the diluted reference standards' concentration using the following equation:

Record the measured flow rates and the undiluted and diluted reference standard concentrations in the laboratory's records.

5. In succession, measure the zero gas, the diluted reference standard(s) and the diluted candidate standard using the analyzer. For each measurement, adjust the flow rates, if necessary, to those determined in step 4, and allow ample time for the analyzer to achieve a stable reading. If the reading for each measurement is not stable, the precision of the measurements will decline and the candidate standard might not be certifiable under this protocol. Record the analyzer response for each measurement, using the same response units (e.g., volts, millivolts, percent of scale, etc.) as was used for the multipoint calibration. At this point, do not convert the data into concentration values using the calibration equation. Do not perform any mathematical transformations of the data. These steps will be done later. Do not make any zero control, span control, or other adjustments to the analyzer during this set of measurements. Record these analyzer responses in the laboratory's records.

The analyst may assay multiple candidate standards during the same assay session. For example, a single set of measurements may involve a zero gas, a diluted reference standard, and three diluted candidate standards. Criteria that apply to the assay of one candidate standard apply to the assay of multiple candidate standards. The analyst should be aware that the effect of any short-term calibration drift will be greater when multiple candidate standards are assayed. This greater effect is due to the longer period of time between reference standard measurements. Unacceptable uncertainties of the estimated concentrations for the diluted candidate standards may occur as a result of the longer assay session.

- 6. Conduct at least two additional sets of measurements, as described in step 5 above. However, for these subsequent sets of measurements, change the order of the three measurements (e.g., measure the reference standard, zero gas, and candidate standard for the second set and measure the zero gas, candidate standard, and reference standard for the third set, etc.). Changing the order that the gas mixtures are measured helps the analyst to discover any effect that one measurement has on subsequent measurements. The number of sets of measurements will have been determined during the analysis of the multipoint calibration data such that the 95-percent uncertainty for the regression-predicted concentration of the candidate standard is ≤1 percent of the concentration of the reference standard.
- 7. If any one or more of the measurements of a set of measurements is invalid or abnormal for any reason, discard all three measurements and repeat the set of measurements. Such measurements may be discarded if the analyst can demonstrate that the experimental conditions were inappropriate during these measurements. Data cannot be discarded just because they appear to be outliers, but may be discarded if they satisfy statistical criteria for testing outliers. The analyst must record the discarded data and a brief explanation as to why these data were discarded in the laboratory's records.
- 8. The spreadsheet described in Appendix A or equivalent statistical techniques must be used to calculate an estimated concentration and a 95-percent uncertainty for the candidate standard based on data from the assay measurements and from the multipoint calibration. The use of both sets of data in the statistical analysis produces an estimated concentration with smaller uncertainty while correcting for any minor calibration drift that may have occurred since the multipoint calibration. Record the estimated concentration and the 95-percent uncertainty in the labora-tory's records.

The spreadsheet also calculates the percentage of the uncertainty that is due to the multipoint calibration. This percentage is needed for the total uncertainty calcula-

tions when two or more assays fall under the same multipoint calibration. Record this value in the laboratory's records.

The analyst should investigate any of the measurements that appear to be outliers. Such data may be discarded if the analyst can demonstrate that the experimental conditions were inappropriate during these measurements. Data cannot be dis-carded just because they appear to be outliers, but may be discarded if they satisfy statistical criteria for testing outliers. The analyst must record any discarded data as well as a brief summary of the investigation in the laboratory's records.

- 9. If the multipoint calibration data and the assay data underwent any mathematical transformations before their statistical analysis, perform the reverse transformations for the estimated concentration and the 95-percent uncertainty. Record the trans-formed values in the laboratory's records.
- 10. Finally, the certified undiluted concentration for a candidate standard containing a unreactive gas mixture and requiring only a single assay can be calculated from the mean concentration of the diluted candidate standard as follows:

where Total Gas Flow Rate = Standard Flow Rate + Zero Gas Flow Rate .

2.3.10 Stability Test for Newly Prepared Standards

Newly prepared candidate standards that contain reactive gas mixtures must be assayed on at least two dates that are separated by at least 7 days. See Subsections 2.1.6.1 and 2.1.6.2. The spreadsheet described in Appendix C or equivalent statistical techniques must be used to evaluate the stability of the candidate standard and to calculate the overall estimated concentration and the total uncertainty for the candidate standard.

The stability is evaluated by comparing the 95-percent confidence limits (i.e., estimated concentration ±95-percent uncertainty) for the candidate standard from the two or more assays. If the confidence limits overlap, the candidate standard can be considered to be stable and may be certified. In the spreadsheet, all cells in the comparisons table will be "true." If the confidence limits do not overlap, the candidate standard may be unstable or there may be analytical problems associated with the assays or the reference standards. One or more cells in the comparisons table will be "false." The analyst must either disqualify the candidate standard or investigate why the confidence limits do not overlap. The analyst may conduct additional assays until stability is achieved and add the additional data to the spreadsheet. Data from a nonoverlapping assay may be discarded and the remaining data used to determine the overall estimated concentration and the total uncertainty provided the confidence limits overlap. Finally calculate a certified undiluted concentration using the above equation. Record these values and any discarded data in the laboratory's records.

2.3.11 Certification Documentation

See Subsections 2.1.4 and 2.1.5.

2.3.12 Recertification Requirements

See Subsections 2.1.6.3 and 2.1.6.4.

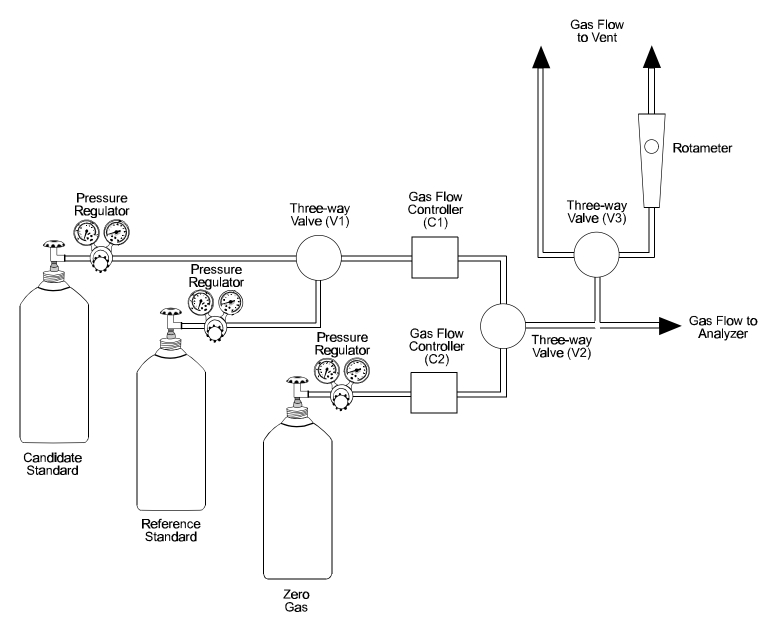


Figure 2-2. One possible design of the apparatus for the assay of compressed gas calibration standards without dilution (Procedure G1)

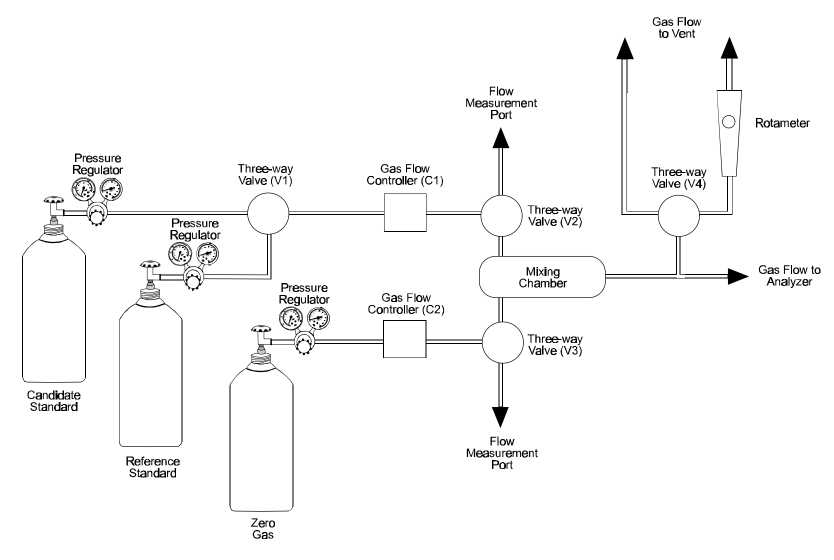


Figure 2-3. One possible design of the apparatus using flow controllers for assay of compressed gas calibration standards with dilution (Procedure G2)

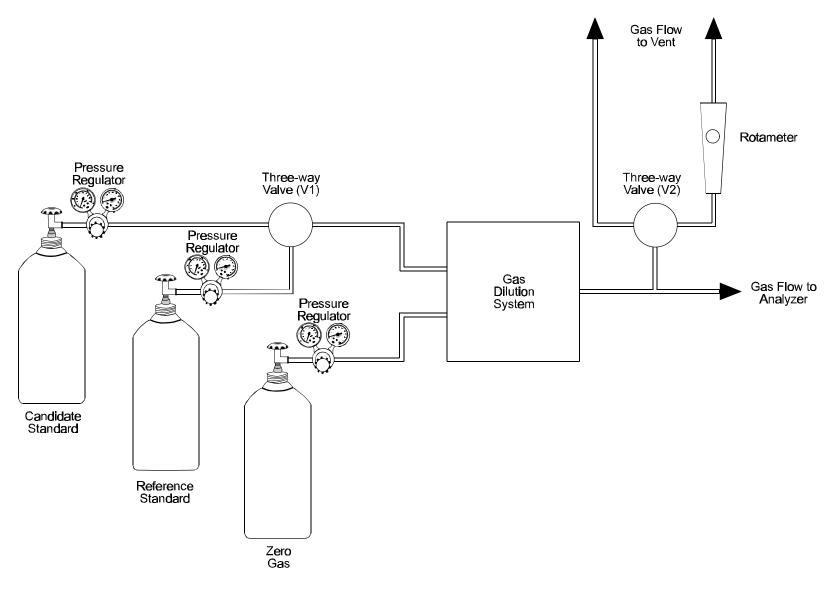


Figure 2-4. One possible design of the apparatus using a gas dilution system for assay of compressed gas calibration standards with dilution (Procedure G2)